

Figure 1

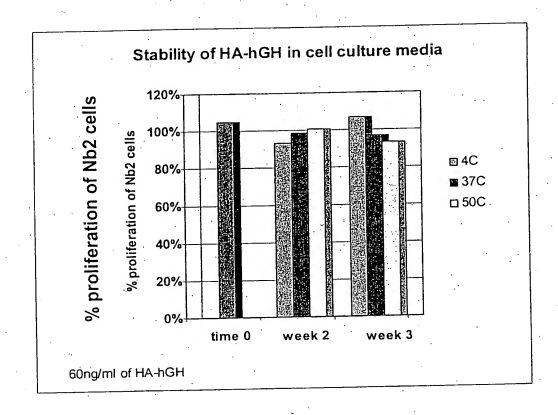


Figure 2

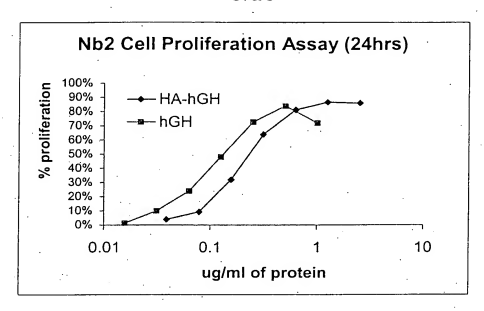


Figure 3A

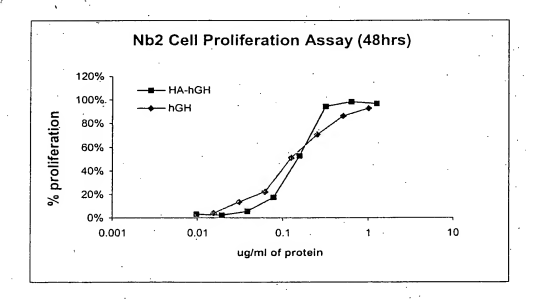


Figure 3B

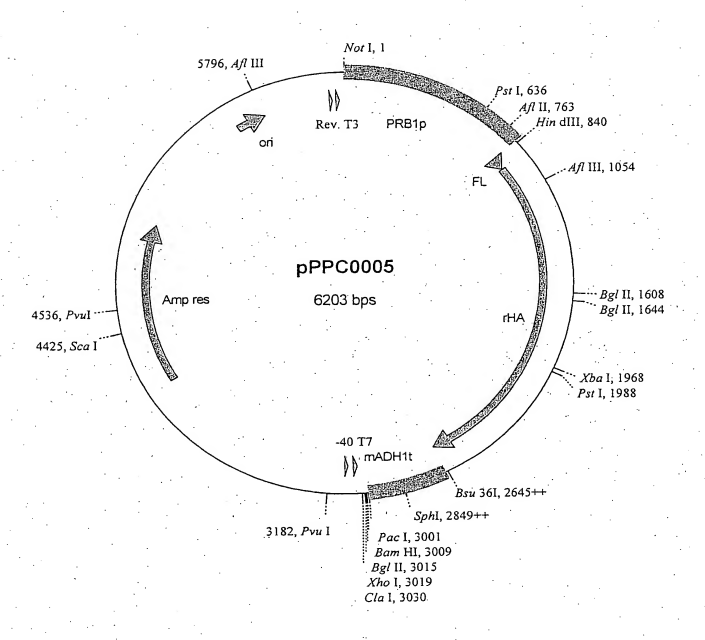


Figure 4

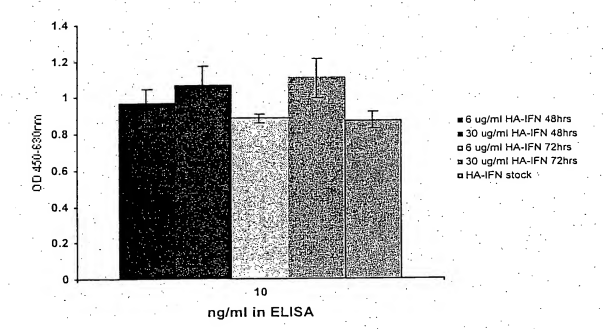
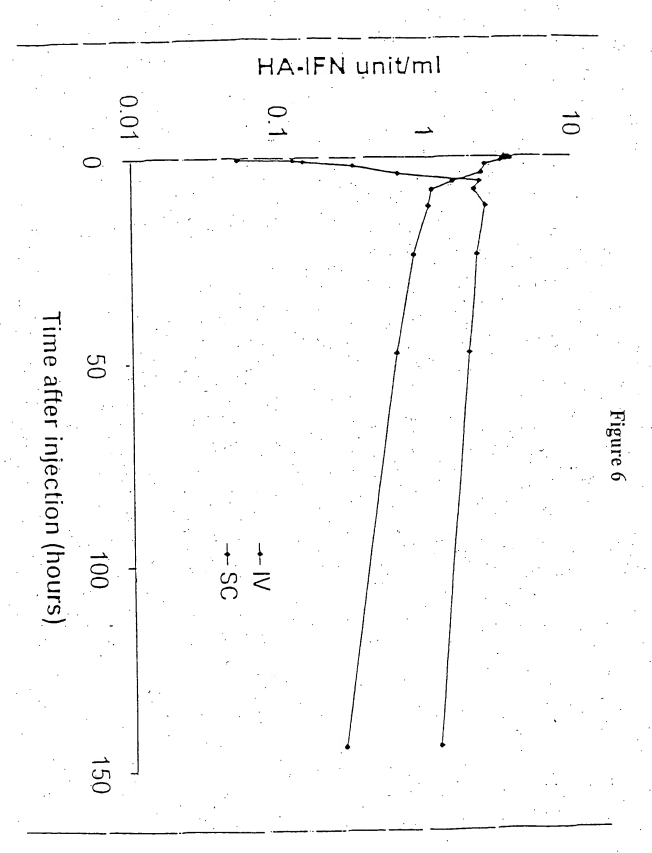
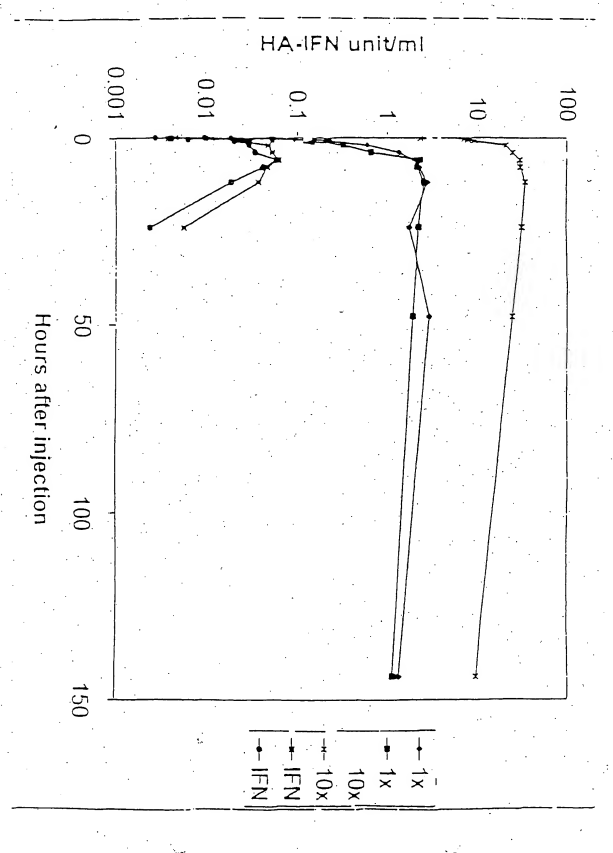


Figure 5





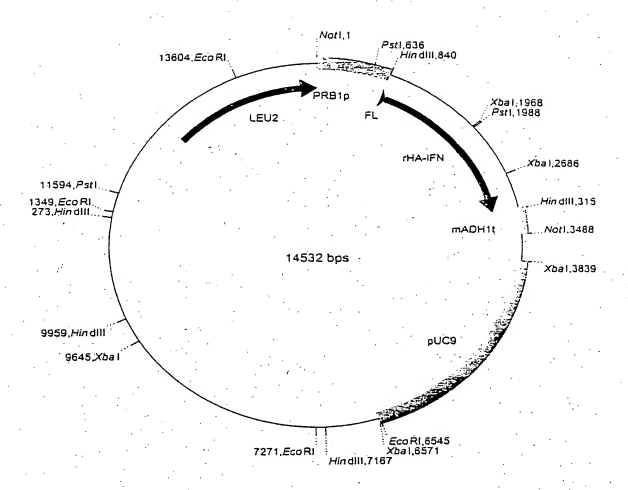


Figure 8. The HA-IFN α expression cassette in pSAC35. The expression cassette comprises

PRB1 promoter, from S. cerevisiae.

Fusion leader, first 19 amino acids of the HA leader followed by the last 6 amino acids of the MF α -1 leader.

HA-IFNa coding sequence with a double stop codon (TAATAA) ADH1 terminator, from S. cerevisiae. Modified to remove all the coding sequence normaly present in the Hind III/BamHI fragment generally used.

Figure 8

9/18

Localisation of 'Loops' based on the HA Crystal Structure which could be used for Mutation/Insertion

1			ALVLІAFAQY НИННИННИН	LQQCPFEDHV ННННН	KLVNEVTEFA ННИНИННИН
	_			<u>.</u>	
	I	• •	•	II	III
51 ·	KTCVADESAE				
	ННННН	НННН	ннннн	. нннн	H HHHH
			•	•	
101	CFLQHKDDNP	NLPRLVRPEV	DVMCTAFHDN	EETFLKKYLY	EIARRHPYFY
	нннн	H	ннннннн	нинининн	нннн
	• • • • • • • • • • • • • • • • • • • •				
IV					
151	APELLFFAKR	YKAAFTECCO	AADKAACLLP	KLDELRDEGK	ASSAKORLKC
				НННЕННННН	
					v
201	ASLOKFGERA	EKAMANADIC	ODEDKAREAE	VCVI VTDI TV	•
•	нинин ни	нинининин	нн ннн	нннннннн	нннннн нн
			_		
			I ·	VII	
2.51		· · —————		KPLLEKSHCI	
	ннннннннн	ННННН	ННННН	нннннн	H
			•		
301	DLPSLAADFV	ESKDVCKNYA	EAKDVFLGMF	LYEYARRHPD	YSVVLLLRLA
	нннн	ннннн	нннннн	ННННН	ННННННН
		•	•••		
	•	. VIII			
351	KTYETTLEKC	CAAADPHECY	AKVEDEEKPL	VEEPONLIKO.	NCELFEOLGE
	ннниннннн	. 		ннннннннн	
					ïx
401	YKFQNALLVR	VTVVIDOUCT	DTT VEVEDNI	CKICCKCCKA	
40T					
	нинининни	пппп п	ппппппппппппппппппппппппппппппппппппппп	nnn	ннннннн
					•
		X		ΧI	•
451	-				LEVDETYVPK
	нннннннн	ННННН	нннннннн	ннннннн	
501	EFNAETFTFH	ADICTLSEKE	RQIKKQTALV	ELVKHKPKAT	KEQLKAVMDD
		ннн ннн	нннименнн	ННН	ннннннн
•	•				
		XII	•		•
551	FAAFVEKCCK	ADDKETCFAE	EGKKLVAASO	AALGL	
	нининин		ннннннннн		
	•				ė
	Loop		Loop		
	-	4 - Aenel	VII	Glu280-His2	
	I Val54-Asn61 II Thr76-Asp89				
			VIII	Ala362-Glu368	
	III Ala92-Glu100		IX	Lys439-Pro447	
	IV Gln170-Ala176		X .	Val462-Lys475	
		47-Glu252	ΧΙ	Thr478-Pro4	
•	VI Glu26	66-Glu277	XII	Lys560-Thr5	66

Figure 9

Examples of Modifications to Loop IV

a. Randomisation of Loop IV.

τv

ΙV

X represents the mutation of the natural amino acid to any other amino acid. One, more or all of the amino acids can be changed in this manner. This figure indicates all the residues have been changed.

b. Insertion (or replacement) of Randomised sequence into Loop IV.

(X)_n

ΙV

The insertion can be at any point on the loop and the length a length where n would typically be 6, 8, 12, 20 or 25.

Figure 10

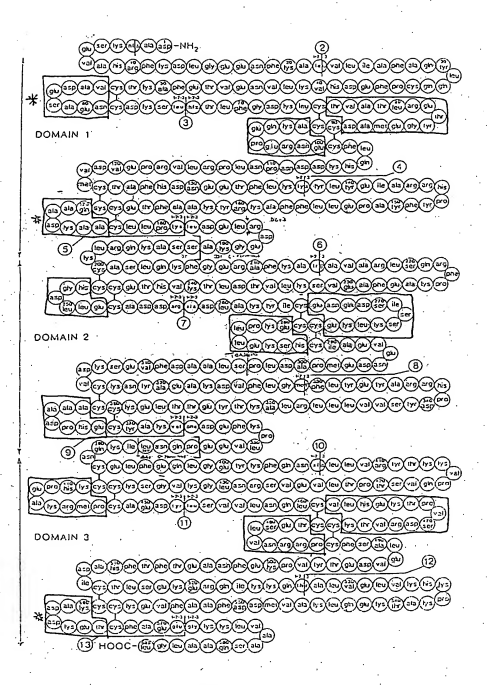
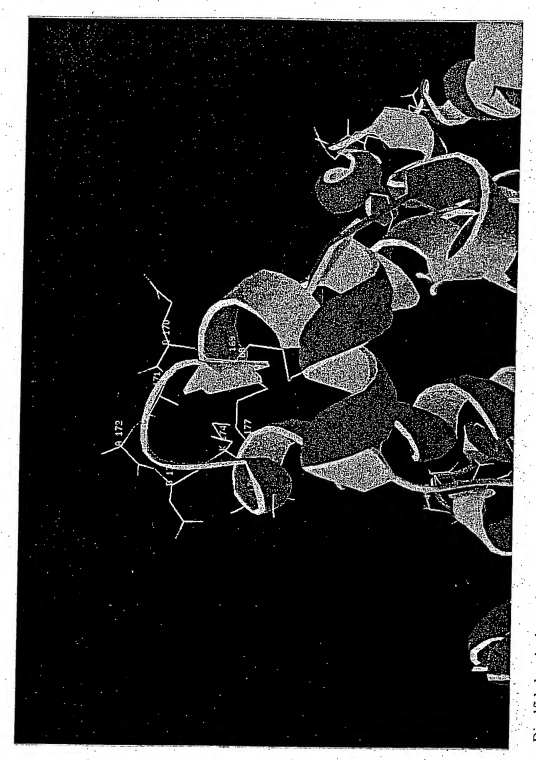


Figure 11



Disulfide bonds shown in yellow

Figure 12: Loop IV Gln170-Ala176

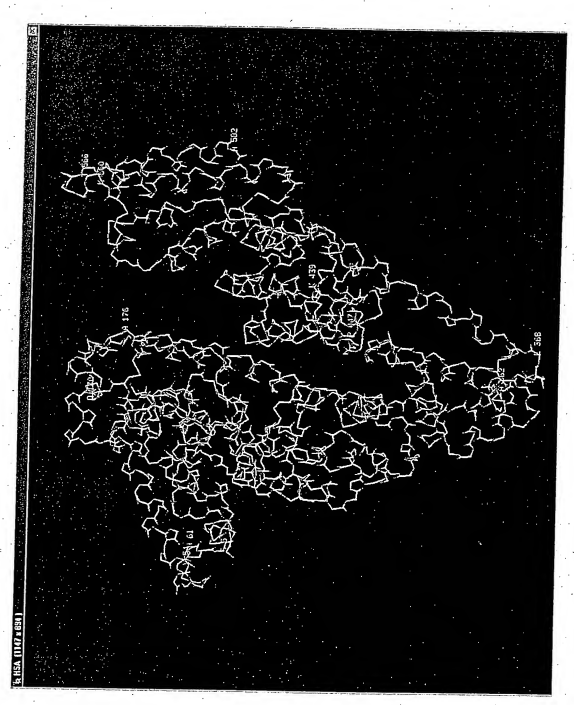


Figure 13: Tertiary Structure of HA

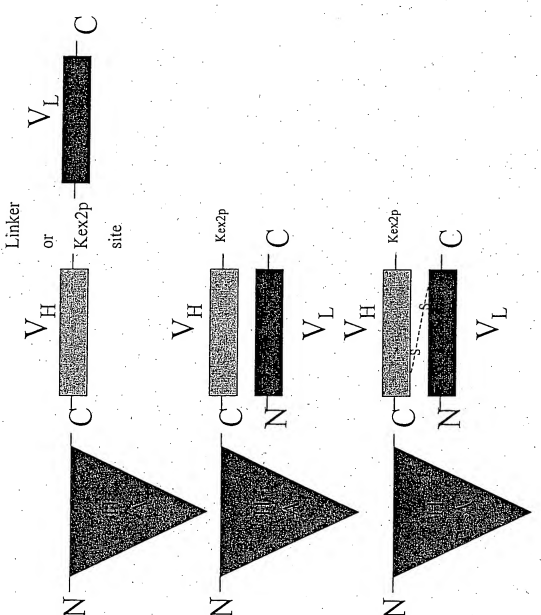


Figure 14: Schematic Diagram of Possible ScFv Fusions (Example is of a C-terminal fusion to HA)

GGA AAG AGT GAT GCT CAT CGG TTT AAA GAT K S E V A H R F K D

120 CCA TTT GAA C TGT CAG CAG 1 TTT GCT CAG TAT CTT F A Q Y L ATT GCC I TTG GTG TTG 61

GAA E GCT GAT GAG TCA D E S GCT ACA TGT GTT (TTT GCA AAA 1 F A K 1 GAA E TTA GTG AAT GAA GTA ACT L V N E V T AAA 121

CTTTGC ACA GTT GCA ACT C T V A T TTA ' GAC AAA O ACC CTT TTT GGA T L F G TGT GAC AAA TCA CTT CAT C D K S L H AAT 181 61

300 GAA E AAT GAA CCT GAG AGA E P E R CAA O GCA AAA O GAC TGC TGT D C C GCT GAA ATG GAA ACC TAT GGT E T Y G 241

360 GAG GTT E. V TGC TTC TTG CAA CAC AAA GAT GAC AAC CCA AAC CTC CCC CGA TTG GTG AGA CCA C F L Q H K D D N P N L P R L V R P 301

420 140 TTA TAT TAC TTG AAA AAA L [Ŧ GAA GAG ACA TTT CAT GAC AAT F H D N GCT Ø TGC ACT ۲ AŢG M GTG V

480 160 AGG R AAA CTC CTT TTC TTT GCT GAA ATT GCC AGA AGA CAT CCT TAC TTT TAT GCC CCG GAA E I A R R H P Y F Y A P E

Figure 15A

TTG CCA E TGCTGT TGC CAA GCT GCT GAT AAA GCT C C Q A A D K A GCT TTT ACA GAA A F T E TAT AAA GCT C Y K A A 161

600 200 $^{\mathrm{TGT}}$ AAA K CTCGCC AAA CAG AGA A K Q R TCG TCT (S S) GCT A AAG K ງ ອີວິ GAA GAT Д CGG R CTT L GAA. E GAT D CTC L AAG . K 541 181

6.60 220 CGC CTG AGC R L S GCT TTC AAA GCA TGG GCA GTG F K A W A V GCT AGA GAA E GGA TTT F AAA K CAA O CTC L AGT S GCC 601 201

720 AAA K ACC T CTT L GAT ACA T TCC AAG-TTA GTG S K L V GTT GAA E TTT CCC AAA GCT GAG TTT GCA F P K A E F A AGA R 661 CAG /

780 260 CTT GAA TGT GCT GAT GAC AGG GCG GAC CTT L E C A D D R A D L CTG L GAT D GGA Ö CAT TGC TGC GAA CAC ACG H T 721 GTC (241 V 1

840 280 GAA TGT C TGC CTG AAG GAA L K E AAA K TCC AGT GAT TCG ATC TD S I 9 CAG ø TAT ATC TGT GAA AAT Y I C E N AAG . ¥ 781 GCC Ø 261

900 300 GAA AAT GAT GAG ATG CCT GCT E N D E M P A GAA GTG E V GCC CAC TGC ATT H C I AAA TCC (TTG GAA I CTG CCT P AAA K 841 281

960 320 TAT GCT Y A AAC TGC AAA GAA AGT AAG GAT GTT E S K D V GTTTCA TTA GCT GCT GAT TTT S L A A D F CCT TTG 901 GAC 3

Figure 15B

CCT GAT 1020 TAT GCA AGA AGG CAT Y A R R H GTC TTC CTG GGC ATG TTT TTG TAT V F L G M F L Y 961 GAG GCA AAG GAT 321 E A K D

AAG CTA GAG GAA ACC ACT E T T GCC AAG ACA TAT A K T Y CTG CTG CTG AGA CTT L L L R L GTC GTG TCT 1021 TAC 341 Y CCT CTT 1140 P L 380 AAA AAA GTG TTC GAT GAA TTT K V F D E F TGC TAT GCC A CCT CAT GAA GCA GAT (1081 TGT GCC GCT 361 C A A

GAG 1200 E 400 GGA CTTGTG GAA GAG CCT CAG AAT TTA ATC AAA CAA AAC TGT GAG CTT TTT GAG CAG V E E P Q N L I K Q N C E L F E Q 1141 381

TTA GTT CGT TAC ACC AAG AAA GTA CCC CAA GTG TCA ACT 1260 L ν R Y T K K ν P Q ν S T 420 TTC CAG AAT GCG CTA . 1 ŏ TAC AAA

AAA CAT 1320 TGT TGT GGC AGC AAA G S K CTT GTA GAG GTC TCA AGA AAC CTA GGA AAA GTG L V E V S R N L G K V ့တ -CCA ACT ். ப் 421 GCA AAA AGA ATG CCC TGT GCA GAA GAC TAT CTA TCC GTG GTC CTG AAC CAG TTA 1380 A K R M P C A B D Y L S V V L N Q L 460 GAA 1321 CCT

GTC ACA AAA TGC TGC ACA GAG TCC 1440 GTG TTG CAT GAG AAA ACG CCA GTA AGT GAC AGA V L H E K T P V S D R $_{
m TGT}$ 1381

Figure 15C

CCC AAA 1441 TTG GTG AAC AGG CGA CCA TGC TTT TCA GCT CTG GAA GTC GAT GAA ACA TAC GTT 481 L V N R R P C F S A L B V D F T V V

ATA TGC ACA CTT TCT GAG AAG GAG 1560 1501 GAG TTT AAT GCT GAA ACA TTC ACC TTC CAT GCA GAT 501 E F N A E T F T F H A D

GAG CTT GTG AAA CAC AAG CCC AAG GCA ACA 1620 E L V K H K P K A T 540 AGA CAA ATC AAG AAA CAA ACT GCA CTT GTT R Q I K K Q T A L V 1561 GCA GCT TTT GTA GAG AAG TGC TGC AAG 1680 A A F V E K C C K 560 GCT GTT ATG GAT GAT TTC Q · Q CTG AAA C 1621 AAA GAG CAA 541 K E Q 1681 GCT GAC GAT AAG GAG ACC TGC TTT GCC GAG GAG GGT AAA AAA CTT GTT GCT GCA AGT CAA 1740 561 A D D K E T C F A E E G K K L V A A S O 580

1741 GCT GCC TTA GGC TTA TAA CAT CTA CAT TTA AAA GCA TCT CAG 1782 581 A A L G L \star

Figure 15D